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(54) Title: LAUNDRY COMPOSITIONS

(57) Abstract

A composition for use in the rinsing of laundry contains particles with perfume located or absorbed in or on the particles, characterised in that the particles are formed of cross-linked organic polymeric material that have a mean particle size not greater than 1 micron. The composition is preferably a fabric softener. A method of treating laundry with the composition is also provided.

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LAUNDRY COMPOSITIONS

This invention relates to the perfuming of rinse conditioner compositions.

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Rinse conditioners are products which are designed to be added to water used for the rinsing of laundry after washing with a detergent composition. Such conditioners contain a material whose function is to confer a benefit to 10 the laundry after the laundry has been rinsed and dried. One of the main benefits delivered by such products is softness.

It is normal to include perfume in such rinse conditioners, 15 firstly to enhance the attractiveness of the product to a user, and secondly to deliver the perfume to the laundry itself.

It is known to incorporate perfume into a porous carrier to 20 reduce the rate at which the perfume evaporates and hence extend the time during which the perfume fragrance remains perceptible.

US-A-4446032 discloses rinse conditioner compositions which 25 contain perfume trapped within carrier particles which may be porous or alternatively may be closed capsules with an impervious layer surrounding the perfume. This document teaches that a suspending agent should be included in the composition.

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JP-A-63/122796 discloses the addition of a polymer latex which contains cationic or tertiary amino groups to a

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detergent composition, for the purpose of enhancing perfume deposition.

WO98/28396 (Quest) discloses perfume absorbed within 5 organic polymer particles and which have a further polymer at their exterior. The polymer particles disclosed desirably have an average particle size of at least 10 µm.

WO 98/28339 (Quest) discloses polymer particles comprising 10 a hydrophobic organic matrix and located at the exterior free cationic groups and a further polymer which comprises free hydroxy groups. The polymer particles disclosed desirably have an average particle size of at least 10 µm.

- 15 EP 397 246 (Minnesota Mining) discloses perfume particles comprising perfume dispersed within certain water-insoluble polymeric materials and encapsulated in a protective shell by coating with a friable coating material.
- 20 EP 617 051 (Allied Colloids) discloses that the controlled release of a fragrance can be provided by incorporating it during emulsion polymerisation of water insoluble monomeric material in which it is dissolved, and by controlling the monomer blend composition and the Tg of the resultant 25 polymer.

The present invention provides a composition for use in the rinsing of laundry, containing particles with perfume located or absorbed in or on the particles, wherein the 30 particles are formed of cross-linked organic polymeric material and have a mean particle size not greater than 1 micron.

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Preferably the polymeric material is a latex.

We have found that cross-linked latices give superior delivery of perfume to fabrics and better physical 5 stability to the product

In preferred forms, this invention is concerned with rinse conditioner compositions which incorporate a material effective to soften fabrics.

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Thus, according to an aspect of this invention there is provided a rinse conditioner composition containing a material effective to soften fabrics, and particles with perfume located or absorbed in or on the particles,

15 characterised in that the particles are formed of crosslinked organic polymeric material, and have a mean particle size not greater than 1 micron.

Incorporating perfume in this way can lead to an increase 20 in the amount of perfume which remains with the fabric after drying, especially when the fabric is dried in a heated tumble dryer.

Additionally particles of this small, colloidal size remain 25 suspended in an aqueous liquid and there is no need to incorporate a separate suspending agent.

In further aspect this invention provides the use of particles of cross-linked organic polymeric material with a 30 mean particle size of not greater than 1 micron and with perfume absorbed therein, in a rinse conditioner.

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A method of treating laundry comprising contacting said laundry with a composition as herein described is also provided.

5 Furthermore, this invention provides a method of making a rinse conditioner which comprises mixing (i) a fabric softening material and (ii) particles of cross-linked organic polymeric material with a mean particle size not greater than 1 micron and with perfume located or absorbed 10 therein or thereon.

In this aspect, the invention can provide a way to incorporate perfume into a rinse conditioner composition while avoiding or mitigating adverse effects on viscosity.

- 15 We have found that perfume can produce adverse changes to the viscosity of rinse conditioner compositions, especially when these contain a substantial percentage of fabric softener. In extreme cases the addition of perfume to a concentrated rinse conditioner composition can cause it to 20 gel and become immobile.
 - Incorporation of perfume in a latex can avoid or mitigate this. This feature of the invention has been found to be applicable in particular to compositions which contain at
- 25 least 8%, more especially at least 12% by weight of fabric softening material. The amount of fabric softening material may lie in a range from 12% to 40% by weight or may be even greater up to 80% or 90% by weight of the composition.
- 30 It is here envisaged in particular that the fabric softening material may be cationic fabric softening material in an amount which is at least half of the total

quantity of fabric softening material present and in addition is at least 8% by weight of the composition, especially at least 12% by weight of the composition.

5 In certain preferred forms, this invention is particularly concerned with cross-linked polymer lactices whose particles have the perfume trapped therein. We have found that cross-linked latices give superior delivery of perfume to fabrics and better physical stability to the product.

10

Therefore, in a further aspect, this invention provides a product for use in the rinsing of laundry, containing perfume trapped within particles of an organic cross-linked polymeric latex, whose particles have a mean particle size 15 not greater than 1 micron.

Compositions in which this invention may be applied and materials used, will now be discussed in more detail.

20 Fabric Softening Rinse Conditioners

A variety of materials are effective to soften fabrics and can be delivered to fabric from an aqueous composition.

A fabric-softening agent functions to give fabric a softer 25 handle. Frequently such agents also provide an anti-static benefit.

Fabric softening agents used in rinse conditioner compositions are usually materials with low solubility in 30 water. Typically the solubility in acidified water at pH 2.5 and 20°C is less than 10g/litre, preferably less than 1g/litre. When added to rinse water such materials form a

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dispersed phase which is then able to deposit on fabrics which are being rinsed in the water.

Many fabric softening materials fall within the categories 5 of amphoteric, zwitterionic, cationic and nonionic compounds.

Amphoteric, zwitterionic and cationic fabric softening

materials include a polar group and usually one or more

10 hydrophobic aliphatic chains such as alkyl or alkenyl
groups each of which contains at least 6 carbon atoms, e.g.
a range from 6 to 50 carbon atoms in a continuous
hydrocarbon chain or a hydrocarbon chain interrupted by a
hereoatom. In many such materials there are either a single

15 aliphatic chain containing 12 to 50 carbon atoms, or a

15 aliphatic chain containing 12 to 50 carbon atoms, or a plurality of aliphatic chains each containing 6 to 30 carbon atoms. Such aliphatic chains many be interrupted by a heteroatom as for instance in an

linkage. Generally it is preferred that a hydrophobic 25 aliphatic chain does not incorporate more than one interrupting heteroatom.

A nonionic fabric softening material will generally include a polar group and an aliphatic hydrocarbon chain of 6 to 30 30 carbon atoms, optionally interrupted by a heteroatom as mentioned above, although hydrocarbon chains of 6 to 18, 7

carbon atoms especially 6 to 11 carbon atoms are preferably without any interruption.

Cationic fabric softeners are of special commercial
5 importance and are a preferred type of softener according
to the invention. Preferred forms of this invention include
cationic fabric softener, optionally with nonionic,
amphoteric or zwitterionic fabric softener present in
addition. Thus the amount of cationic fabric softening
10 agent may be equal to, or greater than, the amount of any
softening agent or surfactant which is not cationic.

Many commercially important fabric softening agents are organic compounds containing nitrogen, and at least one 15 hydrocarbon chain of 6 to 50 carbon atoms. The nitrogen atom may well be quaternary, but need not be, as for example in amines and imidazolines which protonate to a quaternary form in acidic solution.

20 Some specific instances of fabric softening agents are:

 Acyclic quaternary ammonium compounds. These compounds are of the general formula (I)

30

wherein each Q_1 is a hydrocarbyl group containing from 15 to 22 carbon atoms. Q_2 is a saturated alkyl or hydroxy

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alkyl group containing from 1 to 4 carbon atoms. Q_3 may be as defined for Q_1 or Q_2 or may be phenyl, Q_4 may be defined as for Q_1 or Q_2 , and X^- is an anion preferably selected from halide, acetate, methyl sulphate and ethyl sulphate 5 radicals.

Throughout this discussion of fabric softening agents the expression hydrocarbyl group refers to alkyl or alkenyl groups optionally substituted or interrupted by functional 10 groups such as -OH, -O-, CONH, -COO-, etc.

Representative examples of these quaternary softeners include tallow trimethyl ammonium methyl sulphate or chloride; ditallow dimethyl ammonium chloride; ditallow 15 dimethyl ammonium methyl sulphate; dihexadecyl dimethyl ammonium chloride; di(hydrogenated tallow) dimethyl ammonium methyl sulphate or chloride; di (coconut) dimethyl ammonium chloride dihexadecyl diethyl ammonium chloride; dibehenyl dimethyl ammonium chloride.

20

Preferred among these are ditallow dimethyl ammonium chloride, di (hydrogenated tallow) dimethyl ammonium chloride, di (coconut) dimethyl ammonium chloride and di (coconut) dimethyl ammonium methosulphate.

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Examples of commercially available materials in this class are ARQUAD 2C, ARQUAD 2HT, ARQUAD 2T (all ex Akzo), PRAPAGEN WK, PRAPAGEN WKT, DODIGEN 1829 (all ex Clariant), QUERTON 4BG, QUERTON 442 (all ex Akzo), AMMONYX KP, AMMONYX 30 SKD (all ex Millchem), SYNPROLAM FS (ex Unichema). The product names are believed to be trademarks.

2) Alkoxylated Polyamines Alkoxylated polyamines of general formula (II) are known

5
$$Q_{4} - N^{+} \left(CH_{2} \right)_{n} - N^{+} \right) Q_{11} \qquad (1+m) X^{-} \qquad (II)$$

$$Q_{5} \qquad Q_{5} \qquad M$$

- 10 Each Q₄ is a hydrocarbyl group containing from 10 to 30 carbon atoms. The Q₅ groups may be the same or different each representing hydrogen, (-C₂H₄O)_pH, (C₃H₆O)_qH, (C₂H₄O)_p, (C₃H₆O)Q,H, and alkyl group containing from 1 to 3 carbon atoms or the group (CH₂)_n, N(Q₅)₂; n and n' are each an
- 15 integer from 2 to 6, m is an integer from 1 to 5 and p, q and (p' + q') may be numbers such that (p + q + p' + q') does not exceed 25. X- is an anion.

Alkoxylated polyamines suitable for use herein include N20 tallowyl, NN'N'-tris (2 hydroxethyl)-1,3- propane diamine
di-hydro chloride; N-cocyl N,N,N',N' pentamethyl-1,3
propane diammonium dichloride or dimethosulphate; N-stearyl
N,N',N' tris (2-hydroxyethyl) N,N'dimethyl-1,3
propanediammonium dimethyl sulphate; N-palmityl N,N',N'tris
25 (3-hydroxyprolyl)-1,3-propanediammonium dihydrobromide; Ntallowyl N-(3 aminopropyl)-1,3-propanediamine
trihydrochloride.

3. Diamido Quaternary Ammonium Salts

30 Diamido quaternary salts of general formula (III) are also known to be useful as fabric softening agents.

 Q_6 is a divalent alkylene group containing from 1 to 3 carbon atoms. Q_1 , Q_2 , Q_5 and X^- are as defined previously. Examples of suitable materials are methylbis

- 10 (tallowamidoethyl) (2-hydroxyethyl) ammonium methyl sulphate and methyl bis (hydrogenated tallowamido ethyl) (2-hydroxyethyl) ammonium methyl sulphate. These materials are available from Goldschmidt under trade names VARISOFT 222 and VARISOFT 110 respectively and under the trade name 15 ACCOSOFT from Stepan.
 - 4. Ester Quaternary Ammonium Salts

A number of ester group containing quaternary ammonium salts, including those disclosed in EP 345842 A2 (Procter & 20 Gamble), EP 239910 (Procter & Gamble) and US 4137180 (Lever) and incorporated herein by reference, are known to be particularly useful as softening materials. These materials can be represented by generic formulae (IV) and (V) below.

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$$Q_{8}$$
 $Q_{7} - N^{+} - CH_{2} - Y - Z - Q_{10}$
 Q_{9}
 X^{-}
(IV)

5

$$Q_2$$
 $Q_2 - N^+ - (CH_2)_n - CH - CH_2 - Z - Q_{10}$
 Q_2
 Z
 Q_{10}

In formula (IV) Q_7 is a hydrocarbyl group containing 1 to 4 10 carbon atoms, Q_8 is $(CH_2)_n-Z-Q_{10}$ where n is an integer from 1 to 4 or $-Q_{10}$. Q_9 is an alkyl or hydroxyalkyl group of 1 to 4 carbon atoms, or is as defined for Q_8 . Q_{10} is a hydrocarbyl group containing from 12 to 22 carbon atoms and Y can be $-CH(OH)-CH_2-$ or Q_5 , as previously defined. Z can 15 be -O-C(O)-O, C(O)-O or -O-C(O)- and X^- is an anion.

In formula (V) the symbols Q_2 , Q_{10} , Z and X^- have the meanings defined previously.

- 20 Examples of suitable materials based on formula IV are methyl bis-[ethyl(tallowyl)]-2-hydroxyethyl ammonium methyl sulphate (or its olelyl equivalent), N,N-di(tallowyl-oxyethyl)-N,N-dimethyl ammonium chloride; N,N-di(2-tallowyloxy-2-oxo-ethyl)-N,N-dimethyl ammonium chloride;
 25 N,N-di(2-tallowyloxyethylcarbonyl oxyethyl)-N,N-dimethyl
- ammonium chloride; N-(2-tallowloxy-2-ethyl)-N-(2-tallowyl oxo-2-oxyethyl)-N, N-dimethyl ammonium chloride; N,N,N-tri(tallowyl-oxyethyl)-N-methyl ammonium chloride; N-(2-tallowyloxy-2-oxyethyl)-N-(tallowyl-N,N-dimethyl)-ammonium
- 30 chloride. Tallowyl may be replaced with cocoyl, palmoyl, lauryl, oleyl, stearyl and palmityl groups. An illustrative

25

example of a formula V material is 1,2-(hardened) ditallowyloxy-3-trimethyl ammonium propane chloride.

Examples of commercially available materials can be 5 obtained under the trade name STEPANTEX VRH 90 (ex Stepan), AKYPOQUAT (ex Kao) and as mixtures of mono and ditallow esters of 2,3-dihydroxy propane trimethyl ammonium chloride (ex Clarient).

10 5. Quaternary Imidazolinium Salts A further class of cationic softener materials is the imidazolinium salts of generic formula (VI).

15
$$CH_3 - CH_2$$
 Q_{11} CH_2 Q_{11} Q_{12} Q_{13} Q_{14} Q_{15} Q_{1

Wherein Q_{11} is a hydrocarbyl group containing from 6 to 24 carbon atoms, G is -N(H)-, or -O-, or NQ_2 , n is an integer between 1 and 4, and Q_7 is as defined above.

Preferred imidazolinium salts include 1-methyl-1- (tallowylamido) ethyl-2tallowyl-4,5 dihydro imidazolinium methosulphate and 1-methyl-1-(palmitoylamido) ethyl-2- octadecyl-4,5-dihydroimidazolinium chloride. Other useful

30 imidazolinium materials are 2-heptadecyl-1-methyl-1-(2 stearylamido) ethyl imidazolinium chloride and 2- lauryl-1-hydroxyethyl-1-olayl imidazolinium chloride. Also suitable

are the imidazolinium fabric softening components of US 4127489 incorporated here by reference.

Representative commercially available materials are VARISOFT 475 (ex Goldschmidt) and REWOQUAT W7500 (ex Rewo).

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A particular aspect of the invention relates to compositions wherein the amount of fabric softening material which is cationic is a majority of the fabric softening material present.

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Primary Secondary and Tertiary Amines
 Primary, secondary and tertiary amines of general formula
 (VII) are useful as softening agents.

15

(VII)

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wherein Q_{11} is a hydrocarbyl group containing form 6 to 24 carbon atoms, Q_{12} is hydrogen or a hydrocarbyl group containing from 1 to 22 carbon atoms and Q_{13} can be

- 25 hydrogen or Q₇. Preferably amines are protonated with hydrochloric acid, orthophosphoric acid or citric acid or any other similar acids for use in fabric conditioning compositions of this invention.
- 30 7. Alkoxylated Amines
 Alkoxylated amines of general formula (VIII) are also useful as components of this invention.

$$Q_1 - N = (CH_2)_2 - N = Q_{15}$$
 (VIII)

5

wherein Q_{14} is $(C_2H_4O)_xH$, Q_{15} is $(C_2H_4O)_yH$ and Q_{16} is $(C_2H_4O)_zH$ and x+y is within the range 2 to 15 and x+y+z is within the range 3 to 15, m can be 0, 1 or 2 and Q_1 is as previously defined.

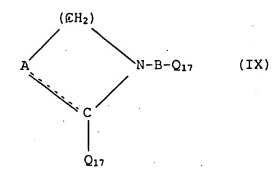
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Examples of these materials are monotallowdipolyethoxyamine containing from 2 to 30 ethylene oxide units, tallow N, N', N' tris (2-hydroxythyl)-1,3 propylene diamine or C_{10} to C_{18} alkyl-N-bis(2-hydroxyethyl) amines.

15 Examples of commercially available materials are available under the trade names ETHOMEEN and ETHODUOMEEN (ex Akzo).

8. Cyclic Amines

Other useful materials are dialkyl cyclic amines 20 represented by formula (IX).



25

wherein the groups Q_{17} are independently selected from hydrocarbyl groups containing from 3 to 30 carbon atoms and A can be oxygen (-O-) or nitrogen (-N-), preferably

15

nitrogen; B is selected from Q_5 as defined earlier or the group $-Q_{13}$ -T-C(O) - where Q_{18} is either Q5 or $(-C_2H_4O_-)_m$ with m being an integer from 1 to 8 and T being selected from oxygen or NQ_{13} . The broken line shows a possible double 5 bond between A and C. If absent, it is replaced by bond(s) to additional hydrogen(s).

Illustrative materials are 12-stearyl oxyethyl-2-stearyl imidazoline, 1-stearyl oxyethyl-2-palmityl imidazoline, 1
10 stearyl oxyethyl myristyl imidazoline, 1-palmityl oxyethyl2-palmityl imidazoline, 1-palmityl oxyethyl-2-myristyl imidazoline, 1-stearyl oxyethyl-2-tallow imidazoline, 1
myristyl oxyethyl-2-tallow imidazoline, 1-palmityl oxyethyl-2-tallow imidazoline, 1-coconut oxyethyl-2-coconut

15 imidazoline, 1-tallow oxyethyl-2-tallow imidazoline and mixtures thereof. Also useful is stearyl hydroxyethyl imidazoline, available commercially as MONAZOLINE S (ex Unichema), 1-tallow amido ethyl-2-tallow imidazoline and

20

Yet another class of suitable fabric softening materials are the condensation products formed from the reaction of fatty acids with a polyamine selected from the group consisting of hydroxyalkyl, alkylene diamines and

Methyl-1-tallow amidoethyl-2-tallow imidazoline.

25 dialkylenetriamines and mixtures thereof. Suitable materials are disclosed in EP-A-199382 (Procter). Preferred among these are mixtures of molecules of the generic formula X and corresponding salts obtained by partial protonation.

25

$$\begin{array}{c} H \\ N - Q_6 - N \\ Q_1 - Q \\ O \end{array}$$

- 10 W is selected from hydrogen and the group $-C(0)-Q_1$ and other symbols are as previously defined. Commercially available materials of this class can be obtained from Clariant as Ceranine HC39, HCA and HCPA.
- 9. Zwitterionic Fabric Softeners
 Other useful ingredients of softening systems include
 zwitterionic quaternary ammonium compounds such as those
 disclosed in EP 332270 A2 (Unilever) incorporated herein by
 reference. Representative materials in this class are
 20 illustrated by general formula (XI) and (XII).

$$Q_{11} - N^{+} - Q_{20} - Z^{-}$$

$$Q_{19}$$
(XI)

30
$$Q_{11} - C - NH - Q_{20} - N^{+} - Q_{20} - Z^{-}$$
 (XII)

wherein the groups Q_{19} are selected independently from Q_7 , Q_{11} and Q_{14} ; Q_{20} is a divalent alkylene group containing 1 to 3 carbon atoms and may be interrupted by -O-, -CONH, - C(0)O-, etc; and Z- is an anionic water solubilising group 5 (e.g. carboxy, sulphate, sulpho or phosphonium).

Examples of commercially available materials are the EMPIGEN CD and BS series (ex Albright & Wilson), the REWOTERIC AM series (ex Goldschmidt) and the Tegobetaine F, 10 H, L and N series (ex GOLDSCHMIDT).

In all of the aforementioned fabric softening compounds where an alkyl chain name is given, the hardened equivalent may also be used, for example tallow or hardened tallow.

15

10. Nonionic Ingredients

It is well known to blend nonionic material with cationic, amphoteric or zwitterionic softening materials as a means of improving dispersion of the product in rinse waters and 20 enhancing the fabric softening properties of the softener blend.

Suitable nonionic adjuncts include lanolin and lanolin derivatives, fatty acids containing from 10 to 18 carbon 25 atoms, esters of fatty acids containing from 8 to 24 carbon atoms with monohydric alcohols containing from 1 to 3 carbon atoms, and polyhydric alcohols containing 2 to 14 carbon atoms eg 2 to 12 carbon atoms such as sucrose, sorbitan, together with alkoxylated fatty acids, alcohols 30 and lanolins containing an average of not more than 7 alkylene oxide groups per molecule. Suitable materials have been disclosed in EP-A-88520 (Unilever), EP-A-122141

(Unilever), GB 2157728A (Unilever), GB 8410321 (Unilever), EP-A-159918 (Unilever), EP-A-159922 (Unilever) and EP-A-79746 (Procter).

- 5 The composition may additionally or alternatively comprise, as a fabric softening agent, an oily sugar derivative which is a liquid or soft solid derivative of a cyclic polyol or of a reduced saccharide, said derivative resulting from 35 to 100% of the hydroxyl groups in said polyol or in said 10 saccharide being esterified or etherified. The derivative has two or more ester or ether groups independently attached to a C_8 - C_{22} alkyl or alkenyl chain. Preferably the oily sugar derivative contains 35% by weight tri or higher
- esters, eg at least 40%. W098/16538 (Unilever) discloses 15 such derivatives and is incorporated herein by reference.

Preferably 35 to 85% most preferably 45 to 70% of the hydroxyl groups in said cyclic polyol or reduced saccharide are esterified or etherified to produce the derivative.

20

The tetra etc prefixes used refer to the average degrees of esterification or etherification; the compounds exist as a mixture comprising the monoester to fully esterified ester.

- 25 The derivatives do not have substantial crystalline character at 20°C. The starting material is esterified or etherified with said alkyl or alkenyl chains to an extent so as to produce the derivatives as a liquid or soft solid.
- 30 Typically the derivative has 3 or more, preferably 4 or more, eg 3 to 5, ester or ether groups or mixtures thereof.

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The derivative may be obtained from a saccharide, e.g. monosaccharides including xylose, arabinose, galactose, fructose, sorbose and especially glucose, or, disaccharides including maltose, lactose, cellobiose and especially 5 sucrose.

An example of a reduced saccharide is sorbitan. If based on a disaccharide it is preferred if the disaccharide has 3 or more ester or ether groups attached to it e.g. include 10 sucrose tri, tetra and penta esters.

Examples of suitable derivatives include esters of alkyl(poly)glucosides, in particular alkyl glucoside esters having a degree of polymerisation from 1 to 2.

15

- The derivative may have branched or linear alkyl or alkenyl chains (of varying degrees of branching), mixed chain lengths and/or unsaturation. Those having unsaturated and/or mixed alkyl chain lengths are preferred.
- 20 For example, predominantly unsaturated fatty chains may be attached to the ester/ether groups, e.g. those derived from rape oil, cotton seed oil, soybean oil, oleic, tallow, palmitoleic, linoleic, erucic or other sources of unsaturated vegetable fatty acids.

25

- Examples of suitable oily sugar derivatives include sucrose tetratallowate, sucrose tetrarapeate, sucrose tetraoleate, sucrose tetraesters of soybean oil or cotton seed oil, cellobiose tetraoleate, sucrose trioleate, sucrose
- 30 triapeate, sucrose pentaoleate, sucrose pentarapeate, sucrose hexaoleate, sucrose hexarapeate, sucrose triesters, pentaesters and hexaesters of soybean oil or cotton seed

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oil, glucose trioleate, glucose tetraoleate, xylose trioleate, sucrose tetralinoleate or sucrose tetra-,tri-, penta- or hexa- esters with any mixture of predominantly unsaturated fatty acid chains.

5

Other oily sugar derivatives suitable include sucrose pentalaurate, sucrose pentaerucate, sucrose tetraerucate. Suitable materials include some of the Ryoto series available from Mitsubishi Kagaku Foods Corporation.

10

The liquid or soft solid derivatives are characterised as materials having a solid:liquid ratio of between 50:50 and 0:100 at 20°C as determined by T_2 relaxation time NMR, preferably between 43:57 and 0:100, most preferably between

- 15 40:60 and 0:100, such as, 20:80 and 0:100. The T_2 NMR relaxation time is commonly used for characterising solid:liquid ratios in soft solid products such as fats and margarines. For the purpose of the present invention, any component of the NMR signal with a T_2 of less than 100
- 20 microsecond is considered to be a solid component and any component with T_2 greater than 100 microseconds is considered to be a liquid component.

The oily sugar derivatives can be prepared by a variety of 25 methods well known to those skilled in the art. Typical preparations of these materials are disclosed in US 4 386 213 and AU 14416/88 (Procter and Gamble).

Fabric softening compositions generally do not contain
30 anionic detergent active nor bleach, nor detergency
builder. It is desirable that the amounts (of any) of
anionic detergent active, bleach and detergency builder are

all less than the amount of the fabric softening agent. The amount of each one of these may well be less than 5% by weight of the whole composition.

- 5 A rinse conditioner composition according to this invention will usually contain from 0.5% to 90% by weight of the fabric softening material, for example 0.5-40%. More specifically, so called dilute compositions generally contain from 1% to about 8% by weight of fabric softening
- 10 agent while compositions containing from about 8% up to 20% by weight of the fabric softening material have been classed as "concentrated". Higher concentrations, in the range of at least 8%, at least 12% or up to 40% by weight are also possible, but even this is not the upper limit.
- 15 Still higher concentrations from 40% up to 80% or even 90% by weight are possible in a very concentrated product.

 As mentioned above, either nonionic or cationic fabric softening agents may be used.

20 Perfume

The term "perfume" as used herein denotes materials which are used in perfumery and mixtures of such materials. Frequently the perfume will be a mixture of perfumery materials. Examples of soitable perfumes are to be found in 25 "Perfume and flavour Chemicals" by Steffen Erctander (Library of Congress Catalogue Card no. 75-91398).

The perfume may function solely to impart fragrance.

However, perfumes can perform other functions, such as to

30 reduce the onset of body malodour as disclosed in US-A
4134838 and EP-A-545556. Such deodorant perfumes are within
the scope of this invention and may or may not impart

22

fragrance. Deodorant perfumes with low perceptible odour are disclosed in EP-A-404470.

Preferably the cross-linked organic polymer particles have 5 the perfume trapped therein.

Polymers

It is a feature of this invention that the particles are composed of cross-linked organic polymeric material. Such

- 10 particles may be made from a variety of organic monomers, although in general these are olefinically unsaturated materials. Preparation may be by an emulsion polymerisation process, as will be explained further below.
- 15 It is also a feature of this invention that the particles have a mean diameter which is not more than lum. Particles of such small size can remain in colloidal suspension. As is known, particles of colloidal dimensions are kept in suspension by Brownian motion and by the effect of charges
- 20 on the particles surfaces, which cause the particles to repel each other and stay apart. Also, because the particles size is small in relation to the wavelength of light, they are generally not visible although they give an appearance of turbidity. Hence, a rinse conditioner
- 25 composition containing them can appear translucent.

There are three preferred techniques for the production of a polymer with perfume absorbed therein as herein described. These are as follows:

30

Method A, is to synthesise a polymer latex by emulsion polymerisation in the presence of the perfume. The monomers

used may include some polyfunctional monomer so as to produce cross-linking in the polymer. This enhances the strength and rigidity of the polymer. A low level of cross-linking provides the ability of particles to hold perfume.

5 Thus, there should desirably be cross linking between at least 0.5%, and 15% by number, of the total number of monomer residues present in the polymer and more preferably between 1% and 10%, eg. between 1 and 5%, of the monomer residues. The polymer particle size range is preferably 10 between 50 and 200 nanometres.

Techniques for the production of a latex of small particles by emulsion polymerisation are well known. The polymerisation is normally carried out in the presence of a 15 surfactant which functions as an emulsifier. Raising the concentration of surfactant leads to smaller particle sizes, but does not generally lead to particle sizes smaller than 50 nanometres unless a special method, e.g. a microemulsion process, is used. Particle size is also 20 affected by temperature and hence rate of reaction, stirring speed and monomer/initiator ratio. Fast rates of polymerisation and higher stirring speeds lead to small particles. High proportions of monomer and low proportions of initiator favour the production of bigger particles.

25

After carrying out emulsion polymerisation it will generally be necessary to remove unreacted monomer. This will often be done by steam distillation, leading to some loss of perfume, although we have found that such losses of perfume are acceptably small. Some monomers may be removed by chemical reaction to yield harmless or volatile products.

Method B: which is particularly preferred, is to commence with a pre-formed cross-linked polymer latex as a dispersion in liquid, e.g. water. The perfume is dissolved in a low boiling organic solvent which is able to swell the polymer particles. The solution is brought into contact with the polymer latex, causing the polymer particles to swell and absorb the perfume. The solvent is then evaporated, leaving the perfume trapped in latex particles.

10 There should desirably be the same amount of cross-linking between of the monomer residues as in method A.

For method B, residual monomer can be removed from the polymer before the polymer comes into contact with the 15 perfume.

For either of these procedures the polymer should be capable of forming a colloidal emulsion in water. A latex of colloidal polymer particles will generally contain from

- 20 3% to 50% polymer, usually from 3% to 40%. Various polymerisation methods may be used for making polymers. Natural or modified natural polymers might also be used, e.g. a natural rubber latex.
- 25 Suitable monomers for radical addition polymerisation are those containing polymerisable olefinic unsaturation.

 Monomers which may be used include styrene, vinyl acetate, ethylene, propylene, vinyl pyrrolidone, vinyl chloride, acylonitrile, methyl methacrylate, and other acrylates and
- 30 methacrylates. The monomers may be used either singly or in combination. Examples of polyfunctional monomers which may be used to effect cross-linking include divinyl benzene,

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divinyl ether, ethyleneglycol dimethacrylate, pentaerythritol triacrylate and polyallyl sucrose. Suitable radical initiators include sodium or potassium persulphate, 2,2'-azobis-isobutyronitrile (AIBN) and 4,4'-azobis (4-5 cyanovaleric acid). A water soluble initiator is preferable. Polymerisation temperature is preferentially in the range 40-90°C.

For Method B, it is possible to use natural and modified 10 natural polymers, e.g. cellulose, modified cellulose, cellulose acetate, cellulose butyrate and hydroxy ethyl cellulose, as well as synthetic latices made from the polymers referred to above.

15 When Method A is used, the perfume should preferably be soluble in the monomer and compatible with the polymer. For Method B, the perfume should also be compatible with the polymer itself. This prevents phase separations which may cause problems.

20 . .

time.

The absorptions of perfume by polymer may be by absorption into open pores through capillary action, or by encapsulation or dissolution in the polymer so that release of perfume is by diffusion through the polymer, or it may 25 be adsorption onto the hydrophobic polymer material. More than one of these modes of absorption may occur at the same

A polymer with absorbed perfume may have the polymer and 30 the perfume in a weight ratio lying in the range from 25:1 to 2:1, especially 20:1 to 3:1.

The amount of polymer particles with absorbed perfume included in a rinse conditioning composition is preferably such an amount that the perfume is present in an amount from 0.1 to 10% by weight of the composition.

5

In many instances, the amount of polymer with absorbed perfume is likely to be such that its perfume amounts to 0.1% to 3%, or even 0.1% to 2% by weight of the composition.

10

A composition according to this invention may incorporate additional perfume which is not located or absorbed in or on the particles (i.e. it is outside the particles and not absorbed thereon) in addition to the perfume which is 15 absorbed onto or into such particles.

References herein to the amount of absorbed perfume in the particles is to be construed as the amount of perfume located in or on the particles, but not perfume separately 20 added to the composition.

If this is done, it can be arranged by choice of the amounts and nature of the perfumes which respectively are and are not absorbed onto the particles that the fragrance 25 on the fabrics after rinsing and drying will come principally from the perfume which was absorbed onto the porous particles whereas the apparent fragrance of the rinse conditioner composition will be principally attributable to the perfume which is not so absorbed. This 30 can enable the producer to control a difference in fragrance between the rinse conditioner composition and the fragrance of the fabrics after rinsing and drying.

27

It may therefore be desirable that the perfume which is not absorbed onto particles, which principally provides the fragrance of the rinse conditioner composition itself, does not exhibit high substantivity towards the fabrics. It may also be desirable that the weight ratio of perfume which is and respectively is not absorbed or included in or on particles lies in a range from 10:1 to 1:4, especially from 10:1 or 4:1 to 1:1.

10

It may be arranged that the polymer melts when heated to ironing temperatures so that perfume which is absorbed onto the polymer particles is released and provides a perceptible fragrance during ironing. If this is desired it 15 will be appropriate for the organic polymer of the porous polymer particles to soften and flow at a temperature in the range from 100°C to 220°C (that is, to have a melting temperature lying in the range 100°C-220°C).

20 Choice of polymer characteristics can affect the ease with which perfume is released. Using monomers which give a more crystalline polymer, and incorporating cross-linking will both inhibit the release of the perfume, so that the perfume may be released more slowly over a longer period or 25 may be released when ironing melts the polymer.

It is preferred that there is cross-linking between at least 0.5% by number of the total number of monomer residues present. There may be cross-linking between up to 30 10% by number of the monomer residues, or possibly even more eg. up to 15% by number. Preferred is a range of 0.5%

or 1% up to 5% or 8% by number of the monomer residues in the polymer.

Other ingredients

- 5 Rinse conditioner compositions may contain a number of materials in addition to the fabric softening material, the porous particles and perfume. Materials which may be present include optical brightening agents, colourants, opacifiers, hydrotropes, viscosity control agents such as
- 10 electrolyte, stabilisers such as guar gum and polyethyleneglycol, anti-static agents and ironing aids. Silicones may be used for the latter purpose.

These various optional ingredients, if present in a rinse 15 conditioner, will generally not be present in amounts exceeding 5% by weight of the composition and they may well total not more than 10% by weight of the composition.

Rinse conditioner compositions in accordance with the
20 invention preferably have a pH of less than 6.0, more
preferably a pH ranging from 2.0 to 5.0. Rinse conditioning
compositions may contain pH-buffering agents such as weak
acids and salts thereof, e.g. phosphoric, benzoic or citric
acids and salts of these acids. The amount of buffering

25 material included in a composition may lie in the range from 0.5-10% by weight, preferably not more than 5% by weight.

Rinse conditioner compositions of this invention are 30 generally in the form of aqueous liquids which will generally contain at least 10%, usually at least 20% by weight water and often from 50% or 60% to 97% by weight of

water. Other product forms may be envisaged, however, including powders, creams, pastes, blocks or tablets. Suitable types of blocks and tablets are disclosed in European Patent Specification No. EP-A-255 779 (UNILEVER). 5 The ability of colloidal particles to remain in suspension is of course most relevant to products in liquid form.

In use compositions according to this invention are preferably added to a much larger volume of water to form a 10 liquor which is than used to contact the fabrics to be treated. Notably, such liquor can be formed by adding a composition by hand or by way of an automatic dispensing device to water in a washing machine.

- 15 One ingredient which is normally not included in rinse conditioning compositions is detergency builder. Generally speaking, rinse conditioning compositions in accordance with this invention will not contain ion exchange aluminosilicate builder, nor any other water-insoluble
- 20 inorganic material nor sodium tripolyphosphate nor any other water-soluble electrolyte in quantities greater than the quantity of the rinse conditioner material. It is likely that the compositions will be entirely free of tripolyphosphate and of any water-insoluble inorganic 25 materials.

The invention will now be further explained and illustrated by the following examples in which all amounts and percentages are by weight unless otherwise indicated.

Example 1

A cross-linked polystrene latex with perfume absorbed therein was prepared by method A referred to above.

- 5 Polymerisation was carried out in a reaction vessel which was a round bottomed flask fitted with stirrer, nitrogen inlet tube and reflux condenser. The materials employed were:
- 10 styrene 8.8 gram
 divinylbenzene (55% concentration) 0.17 gram
 perfume 1 gram
 cetyl trimethylammonium bromide (CTAB) 0.4 gram
 polymerisation initiator 0.25 gram
 water 100 ml.

The CTAB was dissolved in most of the water (90ml) and the solution degassed. The perfume was dissolved in the styrene divinylbenzene mixture and the resulting solution was added

- 20 to the surfactant solution in the flask. After 10 minutes the initiator in the remaining water was added and the polymerisation reaction allowed to proceed for 6-8 hours while the contents of the flask were heated to 70°C.

 Emulsion polymerisation took place during this time giving
- 25 a latex consisting of a colloidal suspension of polymer particles in the water. The mean particle size was approximately 0.1µ. Unreacted monomer was removed by steam distillation. This led to some loss of perfume, but about 90% of the perfume remained in the polymer latex.

A higher quantity of perfume could be incorporated into the latex particles by increasing the quantity of perfume, up to about 2.5 gram.

5 Example 2

In this example perfume was incorporated into performed latex in accordance with method B above.

A latex of cross-linked polystyrene particles of mean size 10 approximately 0.1 μ was prepared using the following materials:

styrene 21.85 gram
divinylbenzene (55%) 4.85 gram

CTAB 0.13 gram
polymerisation initiator 0.33 gram
water 100 gram

The procedure was as given in Example 1 above and after the removal of excess monomer by steam distillation the solids 20 content of the colloidal suspension was determined and found to be 7.8%.

A second stage used:

perfume 1 gram

CTAB 0.3 gram

dichloromethane 27 gram.

The perfume was dissolved in the dichloromethane solvent.

This was then mixed with the CTAB and 50 grams of the latex

30 prepared as above. The heavy organic layer was dispersed throughout the mixture by shaking and then by application

of ulta-sound. This produced an opaque emulsion from which the solvent was removed using a rotary evaporator.

Example 3

5 A typical conditioner composition contains:

wt8.

Dimethyl di-hardened tallow
10 ammonium chloride

5%

Cross-linked Polymer latex
particles according to
any of the preceding examples

sufficient to provide up to 0.4% perfume

15

Demineralised water

balance to 100%

A chemical preservative could well be included, e.g. 0.02% by weight of 2-brom-2-nitropropane-1, 3-diol available 20 under the Registered Trade Mark "Bronopol".

An additional amount of the same perfume, or different, perfume as used in the latex particles can be included in conventional amounts.

25

To provide further examples the dimethyl di-hardened tallow ammonium chloride can be partially, or totally, replaced by 1,2-(hardened) ditallowýloxy-3-trimethyl ammonium propane chloride or a nonionic oily sugar derivative as described 30 in the text above under nonionic compounds.

33

Example 4

Several cross linked polymers latices with perfume incorporated in the latex particles were prepared using the methods A and B disclosed above and exemplified by Examples 1 and 2. Details of these various latices are given in the following Table 1 where percent solids denotes the percentage by weight of polymer in the aqueous latex; percent perfume denotes the quantity of perfume by weight as a percentage of the total weight of polymer and perfume.

10

Each latex was used to deliver perfume to fabric in accordance with the following procedure which used three fabrics namely 70-30 polyester/cotton, a bulked acrylic fabric and cotton terry towelling.

15

Six examples of each of the three fabrics were washed at 50°C using an unperfumed detergent composition containing anionic detergent, phosphate builder and other conventional detergent ingredients.

20

The wash load was rinsed twice with water only and spun dry. At this point the wash load was divided into two halves each containing three pieces of each fabric. One half of the wash load serving as a control was rinsed using 25 a rinse conditioning composition containing the same perfume as that used to make the latices. The other half of the wash load was rinsed using unperfumed rinse conditioning composition mixed with an amount of latex calculated to contain the same quantity of perfume. Each of 30 the halves of the wash load was spun dry, line dried,

stored overnight and then assessed as follows.

The perfume intensity assessment was carried out by a panel of assessors trained to recognise the perfume and score the perfume intensity on a 0-5 scale as extending from 0 = no perfume perceptible through 5 - extremely strong perfume 5 perceived. The panellist's results were averaged and are set out in Table 1 below where " Δ intensity" denotes the amount by which the panellists' mean score exceeded their mean score for the control composition. The results are given in Table 1 below.

10

The two non-cross linked styrene monomer only examples are comparative.

<u>Table 1</u>						
15	Monomers	Method	%Solids	%Perfume	ΔIntensity	Fabric
	STY	В	9.6	20	0.9	Terry
	-				0.9	Polycotton
			•		0.6	Acrylic
20	STY/5% DVB	В	6.5	10	0.7	Terry
					0.8	Acrylic
	STY/10% DVE	B B	5.3	10	1.7	Terry
	•				0.9	Polycotton
25			· .		1.2	Acrylic
	STY	A	12.3	10	0.4	Terry
					0.5	Acrylic

³⁰ From the above results it is demonstrated that the greater the level of DVB, the greater the level of cross-linking and the greater is the perfume substantivity/intensity.

Example 5

Several cross-linked polymer latices with perfume incorporated therein were prepared by the Method B above, exemplified in Example 2. Details are given in the 5 following Table 3.

			Table 2	· .		
	Latex No.	Monomers %	Solids %	Perfume 2	Avge	Particle
10						size (μm)
	I	Styrene/DVB	22	20		0.10
	II	Styrene/DVB	10	20		0.11
	III	Styrene/DVB	23.3	10		0.10

These latices were used to deliver perfume to fabric,

- 15 either terry towelling or polyester cotton mixtures, using a similar procedure to that in the previous Example, or a variant procedure in which the fabric pieces were tumble dried at 70-75°C. The rinse conditioner contained 5% by weight cationic fabric softener (which was
- 20 dimethyldistearyl ammonium chloride). In some experiments the rinse conditioner contained sufficient latex to provide 0.25% by weight perfume. In other experiments the quantity of latex was halved, but compared with a control containing 0.25% by weight perfume. In other experiments the quantity of latex was halved, but compared with a control containing 0.25% by weight of perfume.

Assessment of the dried fabric pieces was carried out as in Example 4 above, except that the results are exprassed as 30 mean perfume scores, and not as Δ relative to a control as in the previous tables.

Details of these experiments and the results obtained are given in the following Table 3.

Table 3

5

Mean Perfume	Intensity	Scores	(0-5	Scale)	On	Dry	Fabric

	•				
		Perfume Level	Fabric	Drying	Mean
	Latex No:	in Composition	Type	Regime	Scores
10	a)			•	
	III	0.25%	Terry	Line	2.2
	II	0.25%	Terry	Line	1.9
	I	0.25%	Terry	Line	2.0
	None (perfumed	0.25%	Terry	Line	1.1
15	control)				
					18
	b)				
•	II	0.25%	Polycotton	Line	1.8
	I	0.25%	Polycotton	Line	1.6
20	None (perfumed	0.25%	Polycotton	Line	1.1
	control)				
				•	
	c)				
	III -	0.125%	Terry	Line	1.2
25	I .	0.125%	Terry	Line	1.2
	None (perfumed	0.25%	Terry	Line	1.2
	control)				
	III	0.125%	Terry	Tumbled	1.8
	I	0.125%	Terry	Tumbled	1.2
30	None (perfumed	0.25%	Terry	Tumbled	1.2
	control)	,			

The results in the above table show that when the perfume levels are equal (at 0.25%) the perfume intensity retained

on the fabric after line drying is greater when the perfume is carried by polymer particles.

The experiments in which the latex provided 0.125% perfume 5 achieved a retention of perfume which was approximately equal to a control using double the quantity of perfume without latex.

Example 6

10 A fabric softening composition was prepared containing 20% by weight of 1, 2-ditallowyloxy-3-trimethyl ammonium propane chloride.

The suspension was a mobile white emulsion.

15

0.5% by weight of perfume was added to this composition. It thickened to such an extent as to become an immobile solid.

The same emulsion was mixed with a latex which was the 20 cross-linked latex I of Example 5. The amount was sufficient to provide 1% by weight perfume in the composition. The emulsion remained very mobile.

The same emulsion was mixed with (1) the same latex in an 25 amount sufficient to provide 0.5% by weight of perfume plus (ii) 0.5% by weight of perfume which was not included in a latex. Again the emulsion thickened dramatically to an immobile solid.

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Example 7

A linear polymeric latex was prepared in the presence of perfume as a comparative example (7A). The materials employed were:

5

	n-butyl acrylate	140g
	N,N'-dimethylaminoethyl acrylate	4.3g
	polyoxyethylene (30) nonylphenyl ether	38.0g
10	perfume	36.1g
	polymerisation initiator	1.2g
	deionised water	500 ml

Polymerisation was carried out in a similar apparatus to 15 that of Example 1. The nonionic surfactant and the polymerisation initiator were dissolved in the water which was then degassed, after which it was stirred and heated to 62°C. The two acrylate monomers were mixed with the perfume and the resulting solution was added drop-wise to the water 20 over a period of two hours. The polymerisation reaction was allowed to continue for a further 7 hours while the contents of the flask were stirred at the same temperature of 62°C. After this the mixture was stem distilled to remove most of the residual monomer. The resulting latex 25 was than concentrated on a rotary evaporator reducing its volume by about 50% and in doing so removing remaining monomer to provide an odourless latex which was filtered through glass wool. 503g of a latex of 37 wt% solids content and 0.11 µm average particle size was obtained.

30

A cross-linked polymeric latex was prepared using the following materials (example 7B):

approximately 0.1u.

	n-butyl acrylate	140 g
	N,N'-dimethylaminoethyl acrylate	4.3 g
٠.	ethylene glycol di(methylacrylate) (EGDMA)	6.0 g
	cetyl trimethyl ammonium bromide (CTAB)	30 g
5	Polymerisation initiator	1.2 g
	deionised water	500 ml
	Polymerisation was carried out in a similar	manner to that
	above. The CTAB was dissolved in the water,	along with the
	initiator. The three acrylate monomers were	mixed together
10	before addition. The EGDMA then functioned a	s a cross-
	linking agent. The latex obtained had a part	icle size of

36.1 grams perfume was incorporated into the resulting
15 latex in accordance with the procedure described in Example
2 above.

The linear and cross-linked latices prepared above
(Examples 7A and 7B respectively) were used to incorporate
20 perfume into a rinse conditioner composition containing 5%
by weight of dimethyl distearyl ammonium chloride as in
Example 6. The amount of each latex was calculated so as to
introduce 0.25% perfume in the rinse conditioner
composition assuming that all perfume used in the
25 preparation of the latex has been retained in the latex.
The resulting rinse conditioner compositions were compared
with a composition containing the same fabric softener and
0.25% by weight perfume without latex.

30 Terry cotton test pieces were agitated in 1 litre of water containing 2 grams of the rinse conditioner under test.

After agitation for 5 minutes the cloths were hand wring,

rinsed in a further litre of plain water for two minutes, then wrung again and line dried overnight. As in Example 4 a panel of assessors scored the perfume intensity on the dried cloths on a scale extending from

 $5 \ 0 = no perfume perceptible through to$

5 = extremely strong perfume perceived

The results of this comparison and a subsequent repetition of it are set out in the following Table 4.

10

TABLE 4 COMPARISON OF LINEAR AND CROSSLINKED POLYMER
LATICES AS PERFUME CARRIERS

SAMPLE	EXPT 1	EXPT 2
Free perfume (control)	0.5	0.7
perfume I cross-linked latex	1.4	1.2
(Example 7B)	*	
perfume in linear latex	1.0	0.9
(Example 7A)		

15 As can be seen from the results above, the perfume incorporated in the cross-linked latex was more perceptible than either of the free perfume control or the perfume incorporated in the linear latex. These results are statistically significant at a 95% confidence level.

20

Using the same test procedure, a comparison was made between latex III of Example 5, the above cross-linked latex based on butyl acrylate (Example 7B) and perfume without latex. As before, the amount of each latex added to 25 the rinse conditioner composition was calculated so as to introduce 0.25% perfume. Some of the test cloths were

ironed before assessment. The results are set out in the following Table 5.

TABLE 5

SAMPLE	EXPT 1	EXPT 1	EXPT 2	EXPT 2
		IRONED		IRONED
free perfume (control	1.0	0.8	0.7	0.6
cross-linked styrene	2.2	3.1	1.6	2.8
latex (Example 5 III)	• .			
cross-linked butyl	1.5	1.3	1.3	1.6 -
acrylate latex		·		
(Example 7B)				

5`

As can be seen from this Table, both cross-linked latices provided an increase in the perceptibility of the perfume delivered to the fabric. This was particularly apparent with the cross-linked styrene latex after the test cloths 10 had been ironed.

Claims

- 1. A composition for use in the rinsing of laundry containing particles with perfume located or absorbed in or on the particles, characterised in that the particles are formed of cross-linked organic polymeric material and have a mean particle size not greater than 1 micron.
- 2. A composition according to claim 1 wherein the polymeric material and the perfume are in a weight ratio lying in the range from 25:1 to 2:1.
- 3. A composition according to claim 1 or claim 2 which is a rinse conditioner composition containing a material effective to soften fabrics.
- 4. A composition according to claim 3 which contains from 0.5 to 90% by weight of the fabric softening material and contains the particles in such amount that the perfume therein or thereon is from 0.1 to 10% by weight of the composition
- 5. A composition according to any preceding claim which is in the form of an aqueous liquid.
- 6. A composition according to claim 4 which contains from 0.5 to 40% by weight of the fabric softening material and contains the particles in such amount that the perfume therein or thereon is from 0.1 to 3% by weight of the composition.

- 7. A composition according to claim 6 which is a liquid composition containing at least 50% by weight of water.
- 8. A composition according to any one of claims 3 to 7 containing at least 12% by weight of the fabric softening material.
- 9. A composition according to any one of claims 3 to 8 wherein the fabric softening material is cationic and is an organic compound containing nitrogen and at least one hydrocarbon chain of 6 to 50 carbon atoms.
- 10. A composition according to any one of claims 3 to 9 containing from 0.5 to 90% by weight of fabric softening material which is cationic.
- 11. A composition according to any one of claims 3 to 10 wherein the amount of fabric softening material which is cationic is a majority of the fabric softening material present.
- 12. A composition according to any one of the preceding claims wherein the particles have the perfume trapped therein.
- 13. A composition according any one of the preceding claims wherein there is cross-linking between at least 0.5% by number of the total number of monomer residues present in the polymer.

- 14. A composition according to any one of the preceding claims which contains additional perfume which is outside the particles, the amounts of perfume which are and are not included in or on the particles lying in a weight ratio from 10:1 to 1:4.
- 15. A composition according to any one of the preceding claims wherein the particles are composed of organic polymer with a melting temperature lying in a range from 100°C to 220°C.
- 16. A method of treating laundry comprising contacting said laundry with a composition as defined in any one of claims 1 to 15.
- 17. A method of making a rinse conditioner comprising mixing;
 - (i) a fabric softening material and
 - (ii) particles of cross-linked organic polymeric material with a mean particle size not greater than 1 micron and with perfume located or absorbed therein of thereon.

INTERNATIONAL SEARCH REPORT

Inter mail Application No PCT/EP 00/03279

			
IPC 7	IFICATION OF SUBJECT MATTER C11D3/50 C11D3/37		•
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According t	o International Patent Classification (IPC) or to both national class	ification and IPC	
	SEARCHED		
Minimum de IPC 7	ocumentation searched (classification system followed by classific C11D	cation symbols)	
	*		
Documenta	tion searched other than minimum documentation to the extent the	at such documents are included in the fields se	arched
Electronic d	lata base consulted during the international search (name of data	base and, where practical, search terms used	
WPI Da	ta, EPO-Internal, PAJ		÷
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to daim No.
X	EP 0 617 051 A (ALLIED COLLOIDS 28 September 1994 (1994-09-28)	LTD)	1-3,12, 16,17
	cited in the application		
A	page 6, line 7 - line 14; claim	l; example	4-11,
			13-15
Α .	WO 98 28396 A (QUEST INTERNATION 2 July 1998 (1998-07-02)	NAL B.V.)	1-17
	cited in the application claims 1,10; examples 6,7		
A	EP 0 397 245 A (PROCTER & GAMBLE	E) *	1-17
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Name and ma	ailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2	Authorized officer	
	NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,		***
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- (22) Date of filing: 10.12.1998
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 MC NL PT SE

 Designated Extension States:

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- (30) Priority: 16.12.1997 EP 97122122
- (71) Applicant: Givaudan SA 1214 Vernier-Genève (CH)
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(51) Int CI.7: **A61K 7/46**, A61K 7/32, A61L 9/04

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- (54) Polymer with binding capacity for organoleptic substances
- (57) The invention relates to a polymer presenting binding sites for at least one organoleptic substance, and specifically, which binding sites are at least partly molecularly imprinted with an organoleptic substance. The polymer which has the property to bind specific classes of organoleptic molecules can be used in any

application where a long lasting effect of the organoleptic substance is targeted, e.g. in cosmetic products, deodorants, air refreshing products, laundry products and in fibers for fabrics. If the polymer prior has been imprinted with malodor molecules this polymer is used to reduce malodor.



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